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(54) Title: INDIVIDUALIZED ANTI-CANCER ANTIBODIES

(57) Abstract: The present invention relates to a method for producing patient specific anti-cancer antibodies using a novel paradigm of screening. By segregating the anti-cancer antibodies using cancer cell cytotoxicity as end point, the process makes possible the production of anti-cancer antibodies customized for the individual patient that can be used for therapeutic and diagnostic purposes.

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INDIVIDUALIZED ANTI-CANCER ANTIBODIES**Reference to Related Applications:**

This application is a continuation-in-part of application S.N. 09/415,278, filed October 8, 1999, now U.S. Patent 6,180,357, the contents of which are herein incorporated by reference.

**Field of the Invention:**

This invention relates to the production of anti-cancer antibodies customized for the individual patient which may be combined with chemotherapeutic agents that can be used for therapeutic and diagnostic purposes. The invention further relates to the process by which the antibodies are made and to their methods of use.

**Background of the Invention:**

Each individual who presents with cancer is unique and has a cancer that is as different from other cancers as that person's identity. Despite this, current therapy treats all patients with the same type of cancer, at the same stage, in the same way. At least 30% of these patients will fail the first line therapy, thus leading to further rounds of treatment and the increased probability of treatment failure, metastases, and ultimately, death. A superior approach to treatment would be the customization of therapy for the particular individual. The only current therapy which lends itself to customization is surgery. Chemotherapy and radiation treatment can not be tailored to the patient, and surgery by itself, in most cases is inadequate for producing cures.

With the advent of monoclonal antibodies, the possibility of developing methods for customized therapy became more realistic since each antibody can be directed to a single epitope. Furthermore, it is possible to produce a combination of antibodies that are directed to

1 the constellation of epitopes that uniquely define a  
2 particular individual's tumor.

3 Having recognized that a significant difference  
4 between cancerous and normal cells is that cancerous cells  
5 contain antigens that are specific to transformed cells,  
6 the scientific community has long held that monoclonal  
7 antibodies can be designed to specifically target  
8 transformed cells by binding specifically to these cancer  
9 antigens; thus giving rise to the belief that monoclonal  
10 antibodies can serve as "Magic Bullets" to eliminate  
11 cancer cells.

12 At the present time, however, the cancer patient  
13 usually has few options of treatment. The regimented  
14 approach to cancer therapy has produced improvements in  
15 global survival and morbidity rates. However, to the  
16 particular individual, these improved statistics do not  
17 necessarily correlate with an improvement in their  
18 personal situation.

19 Thus, if a methodology was put forth which enabled  
20 the practitioner to treat each tumor independently of  
21 other patients in the same cohort, this would permit the  
22 unique approach of tailoring therapy to just that one  
23 person. Such a course of therapy would, ideally, increase  
24 the rate of cures, and produce better outcomes, thereby  
25 satisfying a long-felt need.

26 Historically, the use of polyclonal antibodies has  
27 been used with limited success in the treatment of human  
28 cancers. Lymphomas and leukemias have been treated with  
29 human plasma, but there were few prolonged remission or  
30 responses. Furthermore, there was a lack of  
31 reproducibility and there was no additional benefit  
32 compared to chemotherapy. Solid tumors such as breast  
33 cancers, melanomas and renal cell carcinomas have also  
34 been treated with human blood, chimpanzee serum, human

1 plasma and horse serum with correspondingly unpredictable  
2 and ineffective results.

3 There have been many clinical trials of monoclonal  
4 antibodies for solid tumors. In the 1980s there were at  
5 least four clinical trials for human breast cancer which  
6 produced only one responder from at least 47 patients  
7 using antibodies against specific antigens or based on  
8 tissue selectivity. It was not until 1998 that there was  
9 a successful clinical trial using a humanized anti-her 2  
10 antibody in combination with Cisplatin. In this trial 37  
11 patients were accessed for responses of which about a  
12 quarter had a partial response rate and another half had  
13 minor or stable disease progression.

14 The clinical trials investigating colorectal cancer  
15 involve antibodies against both glycoprotein and  
16 glycolipid targets. Antibodies such as 17-1A, which has  
17 some specificity for adenocarcinomas, had undergone Phase  
18 2 clinical trials in over 60 patients with only one  
19 patient having a partial response. In other trials, use  
20 of 17-1A produced only one complete response and two minor  
21 responses among 52 patients in protocols using additional  
22 cyclophosphamide. Other trials involving 17-1A yielded  
23 results that were similar. The use of a humanized murine  
24 monoclonal antibody initially approved for imaging also  
25 did not produce tumor regression. To date there has not  
26 been an antibody that has been effective for colorectal  
27 cancer. Likewise there have been equally poor results for  
28 lung cancer, brain cancers, ovarian cancers, pancreatic  
29 cancer, prostate cancer, and stomach cancer. There has  
30 been some limited success in the use of anti-GD3  
31 monoclonal antibody for melanoma. Thus, it can be seen  
32 that despite successful small animal studies that are a  
33 prerequisite for human clinical trials, the antibodies  
34 that have been tested have been for the most part  
35 ineffective.

1     **Prior Patents:**

2             U.S. Patent No. 5,750,102 discloses a process wherein  
3     cells from a patient's tumor are transfected with MHC  
4     genes which may be cloned from cells or tissue from the  
5     patient. These transfected cells are then used to  
6     vaccinate the patient.

7             U.S. Patent No. 4,861,581 discloses a process  
8     comprising the steps of obtaining monoclonal antibodies  
9     that are specific to an internal cellular component of  
10    neoplastic and normal cells of the mammal but not to  
11    external components, labeling the monoclonal antibody,  
12    contacting the labeled antibody with tissue of a mammal  
13    that has received therapy to kill neoplastic cells, and  
14    determining the effectiveness of therapy by measuring the  
15    binding of the labeled antibody to the internal cellular  
16    component of the degenerating neoplastic cells. In  
17    preparing antibodies directed to human intracellular  
18    antigens, the patentee recognizes that malignant cells  
19    represent a convenient source of such antigens.

20            U.S. Patent No. 5,171,665 provides a novel antibody  
21    and method for its production. Specifically, the patent  
22    teaches formation of a monoclonal antibody which has the  
23    property of binding strongly to a protein antigen  
24    associated with human tumors, e.g. those of the colon and  
25    lung, while binding to normal cells to a much lesser  
26    degree.

27            U.S. Patent No. 5,484,596 provides a method of cancer  
28    therapy comprising surgically removing tumor tissue from a  
29    human cancer patient, treating the tumor tissue to obtain  
30    tumor cells, irradiating the tumor cells to be viable but  
31    non-tumorigenic, and using these cells to prepare a  
32    vaccine for the patient capable of inhibiting recurrence  
33    of the primary tumor while simultaneously inhibiting  
34    metastases. The patent teaches the development of  
35    monoclonal antibodies which are reactive with surface  
36    antigens of tumor cells. As set forth at col. 4, lines 45

1 et seq., the patentees utilize autochthonous tumor cells  
2 in the development of monoclonal antibodies expressing  
3 active specific immunotherapy in human neoplasia.

4 U.S. Patent No. 5,693,763 teaches a glycoprotein  
5 antigen characteristic of human carcinomas and not  
6 dependent upon the epithelial tissue of origin.

7 U.S. Patent No. 5,783,186 is drawn to Anti-Her2  
8 antibodies which induce apoptosis in Her2 expressing  
9 cells, hybridoma cell lines producing the antibodies,  
10 methods of treating cancer using the antibodies and  
11 pharmaceutical compositions including said antibodies.

12 U.S. Patent No. 5,849,876 describes new hybridoma  
13 cell lines for the production of monoclonal antibodies to  
14 mucin antigens purified from tumor and non-tumor tissue  
15 sources.

16 U.S. Patent No. 5,869,268 is drawn to a method for  
17 producing a human lymphocyte producing an antibody  
18 specific to a desired antigen, a method for producing a  
19 monoclonal antibody, as well as monoclonal antibodies  
20 produced by the method. The patent is particularly drawn  
21 to the production of an anti-HD human monoclonal antibody  
22 useful for the diagnosis and treatment of cancers.

23 U.S. Patent No. 5,869,045 relates to antibodies,  
24 antibody fragments, antibody conjugates and single chain  
25 immunotoxins reactive with human carcinoma cells. The  
26 mechanism by which these antibodies function is two-fold,  
27 in that the molecules are reactive with cell membrane  
28 antigens present on the surface of human carcinomas, and  
29 further in that the antibodies have the ability to  
30 internalize within the carcinoma cells, subsequent to  
31 binding, making them especially useful for forming  
32 antibody-drug and antibody-toxin conjugates. In their  
33 unmodified form the antibodies also manifest cytotoxic  
34 properties at specific concentrations.

35 U.S. Patent No. 5,780,033 discloses the use of  
36 autoantibodies for tumor therapy and prophylaxis. However,

1 this antibody is an antinuclear autoantibody from an aged  
2 mammal. In this case, the autoantibody is said to be one  
3 type of natural antibody found in the immune system.  
4 Because the autoantibody comes from "an aged mammal",  
5 there is no requirement that the autoantibody actually  
6 comes from the patient being treated. In addition the  
7 patent discloses natural and monoclonal antinuclear  
8 autoantibody from an aged mammal, and a hybridoma cell  
9 line producing a monoclonal antinuclear autoantibody.

10

11 **Summary of the Invention:**

12 This application teaches a method for producing  
13 patient specific anti-cancer antibodies using a novel  
14 paradigm of screening. These antibodies can be made  
15 specifically for one tumor and thus make possible the  
16 customization of cancer therapy. Within the context of  
17 this application, anti-cancer antibodies having either  
18 cell-killing (cytotoxic) or cell-growth inhibiting  
19 (cytostatic) properties will hereafter be referred to as  
20 cytotoxic. These antibodies can be used in aid of staging  
21 and diagnosis of a cancer, and can be used to treat tumor  
22 metastases.

23 The prospect of individualized anti-cancer treatment  
24 will bring about a change in the way a patient is managed.  
25 A likely clinical scenario is that a tumor sample is  
26 obtained at the time of presentation, and banked. From  
27 this sample, the tumor can be typed from a panel of pre-  
28 existing anti-cancer antibodies. The patient will be  
29 conventionally staged but the available antibodies can be  
30 of use in further staging the patient. The patient can be  
31 treated immediately with the existing antibodies, and a  
32 panel of antibodies specific to the tumor can be produced  
33 either using the methods outlined herein or through the  
34 use of phage display libraries in conjunction with the  
35 screening methods herein disclosed. All the antibodies  
36 generated will be added to the library of anti-cancer



1     antibodies since there is a possibility that other tumors  
2     can bear some of the same epitopes as the one that is  
3     being treated.

4             In addition to anti-cancer antibodies, the patient  
5     can elect to receive the currently recommended therapies  
6     as part of a multi-modal regimen of treatment. The fact  
7     that the antibodies isolated via the present methodology  
8     are relatively non-toxic to non-cancerous cells allow  
9     combinations of antibodies at high doses to be used,  
10    either alone, or in conjunction with conventional therapy.  
11    The high therapeutic index will also permit re-treatment  
12    on a short time scale that should decrease the likelihood  
13    of emergence of treatment resistant cells.

14            If the patient is refractory to the initial course of  
15    therapy or metastases develop, the process of generating  
16    specific antibodies to the tumor can be repeated for re-  
17    treatment. Furthermore, the anti-cancer antibodies can be  
18    conjugated to red blood cells obtained from that patient  
19    and re-infused for treatment of metastases. There have  
20    been few effective treatments for metastatic cancer and  
21    metastases usually portend a poor outcome resulting in  
22    death. However, metastatic cancers are usually well  
23    vascularized and the delivery of anti-cancer antibodies by  
24    red blood cells can have the effect of concentrating the  
25    antibodies at the site of the tumor. Even prior to  
26    metastases, most cancer cells are dependent on the host's  
27    blood supply for their survival and anti-cancer antibody  
28    conjugated red blood cells can be effective against *in*  
29    *situ* tumors, too. Alternatively, the antibodies may be  
30    conjugated to other hematogenous cells, e.g. lymphocytes,  
31    macrophages, monocytes, natural killer cells, etc.

32  
33            There are five classes of antibodies and each is  
34    associated with a function that is conferred by its heavy  
35    chain. It is generally thought that cancer cell killing

1 by naked antibodies are mediated either through antibody  
2 dependent cellular cytotoxicity or complement dependent  
3 cytotoxicity. For example murine IgM and IgG2a antibodies  
4 can activate human complement by binding the C-1 component  
5 of the complement system thereby activating the classical  
6 pathway of complement activation which can lead to tumor  
7 lysis. For human antibodies the most effective complement  
8 activating antibodies are generally IgM and IgG1. Murine  
9 antibodies of the IgG2a and IgG3 isotype are effective at  
10 recruiting cytotoxic cells that have Fc receptors which  
11 will lead to cell killing by monocytes, macrophages,  
12 granulocytes and certain lymphocytes. Human antibodies of  
13 both the IgG1 and IgG3 isotype mediate ADCC.

14 Another possible mechanism of antibody mediated  
15 cancer killing may be through the use of antibodies that  
16 function to catalyze the hydrolysis of various chemical  
17 bonds in the cell membrane and its associated  
18 glycoproteins or glycolipids, so-called catalytic  
19 antibodies.

20 There are two additional mechanisms of antibody  
21 mediated cancer cell killing which are more widely  
22 accepted. The first is the use of antibodies as a vaccine  
23 to induce the body to produce an immune response against  
24 the putative cancer antigen that resides on the tumor  
25 cell. The second is the use of antibodies to target  
26 growth receptors and interfere with their function or to  
27 down regulate that receptor so that effectively its  
28 function is lost.

29 Accordingly, it is an objective of the invention to  
30 teach a method for producing anti-cancer antibodies from  
31 cells derived from a particular individual which are  
32 cytotoxic with respect to cancer cells while  
33 simultaneously being relatively non-toxic to non-cancerous  
34 cells.

35 It is an additional objective of the invention to  
36 produce novel anti-cancer antibodies.

1           It is a further objective of the instant invention to  
2 produce anti-cancer antibodies whose cytotoxicity is  
3 mediated through antibody dependent cellular toxicity.

4           It is yet an additional objective of the instant  
5 invention to produce anti-cancer antibodies whose  
6 cytotoxicity is mediated through complement dependent  
7 cellular toxicity.

8           It is still a further objective of the instant  
9 invention to produce anti-cancer antibodies whose  
10 cytotoxicity is a function of their ability to catalyze  
11 hydrolysis of cellular chemical bonds.

12           Still an additional objective of the instant  
13 invention is to produce anti-cancer antibodies useful as a  
14 vaccine to produce an immune response against putative  
15 cancer antigen residing on tumor cells.

16           A further objective of the instant invention is the  
17 use of antibodies to target cell membrane proteins, such  
18 as growth receptors, cell membrane pumps and cell  
19 anchoring proteins, thereby interfering with or down  
20 regulating their function.

21           Yet an additional objective of the instant invention  
22 is the production of anti-cancer antibodies whose cell-  
23 killing utility is concomitant with their ability to  
24 effect a conformational change in cellular proteins such  
25 that a signal will be transduced to initiate cell-killing.

26           A still further objective of the instant invention is  
27 to produce anti-cancer antibodies which are useful for  
28 diagnosis, prognosis, and monitoring of cancer, e.g.  
29 production of a panel of therapeutic anti-cancer  
30 antibodies to test patient samples to determine if they  
31 contain any suitable antibodies for therapeutic use.

32           Yet another objective of the instant invention is to  
33 produce novel antigens, associated with cancer processes,  
34 which can be discovered by using anti-cancer antibodies  
35 derived by the process of the instant invention. These

1 antigens are not limited to proteins, as is generally the  
2 case with genomic data; they may also be derived from  
3 carbohydrates or lipids or combinations thereof.

4 Other objects and advantages of this invention will  
5 become apparent from the following description wherein are  
6 set forth, by way of illustration and example, certain  
7 embodiments of this invention.

8

#### 9 Detailed Description of the Invention:

10 It is to be understood that while a certain form of  
11 the invention is illustrated, it is not to be limited to  
12 the specific form or arrangement herein described and  
13 shown. It will be apparent to those skilled in the art  
14 that various changes may be made without departing from  
15 the scope of the invention and the invention is not to be  
16 considered limited to what is shown and described in the  
17 specification.

18 One of the potential benefits of monoclonal  
19 antibodies with respect to the treatment of cancer is  
20 their ability to specifically recognize single antigens.  
21 It was thought that in some instances cancer cells possess  
22 antigens that were specific to that kind of transformed  
23 cell. It is now more frequently believed that cancer  
24 cells have few unique antigens, rather, they tend to over-  
25 express a normal antigen or express fetal antigens.  
26 Nevertheless, the use of monoclonal antibodies provided a  
27 method of delivering reproducible doses of antibodies to  
28 the patient with the expectation of better response rates  
29 than with polyclonal antibodies.

30 Traditionally, monoclonal antibodies have been made  
31 according to fundamental principles laid down by Kohler  
32 and Milstein. Mice are immunized with antigens, with or  
33 without, adjuvants. The splenocytes are harvested from  
34 the spleen for fusion with immortalized hybridoma  
35 partners. These are seeded into microtitre plates where

1 they can secrete antibodies into the supernatant that is  
2 used for cell culture. To select from the hybridomas that  
3 have been plated for the ones that produce antibodies of  
4 interest the hybridoma supernatants are usually tested for  
5 antibody binding to antigens in an ELISA (enzyme linked  
6 immunosorbent assay) assay. The idea is that the wells  
7 that contain the hybridoma of interest will contain  
8 antibodies that will bind most avidly to the test antigen,  
9 usually the immunizing antigen. These wells are then  
10 subcloned in limiting dilution fashion to produce  
11 monoclonal hybridomas. The selection for the clones of  
12 interest is repeated using an ELISA assay to test for  
13 antibody binding. Therefore, the principle that has been  
14 propagated is that in the production of monoclonal  
15 antibodies the hybridomas that produce the most avidly  
16 binding antibodies are the ones that are selected from  
17 among all the hybridomas that were initially produced.  
18 That is to say, the preferred antibody is the one with  
19 highest affinity for the antigen of interest.

20 There have been many modifications of this procedure  
21 such as using whole cells for immunization. In this  
22 method, instead of using purified antigens, entire cells  
23 are used for immunization. Another modification is the  
24 use of cellular ELISA for screening. In this method  
25 instead of using purified antigens as the target in the  
26 ELISA, fixed cells are used. In addition to ELISA tests,  
27 complement mediated cytotoxicity assays have also been  
28 used in the screening process. However, antibody-binding  
29 assays were used in conjunction with cytotoxicity tests.  
30 Thus, despite many modifications, the process of producing  
31 monoclonal antibodies relies on antibody binding to the  
32 test antigen as an endpoint.

33 Most antibodies directed against cancer cells have  
34 been produced using the traditional methods outlined  
35 above. These antibodies have been used both  
36 therapeutically and diagnostically. In general, for both

1 these applications, the antibody has been used as the  
2 targeting agent that delivers a payload to the site of the  
3 cancer. These antibody conjugates can either be  
4 radioactive, toxic, or serve as an intermediary for  
5 further delivery of a drug to the body, such as an enzyme  
6 or biotin. Furthermore, it was widely held, until  
7 recently, that naked antibodies had little effect in vivo.  
8 Both HERCEPTIN and RITUXIMAB are humanized murine  
9 monoclonal antibodies that have recently been approved for  
10 human use by the FDA. However, both these antibodies were  
11 initially made by assaying for antibody binding and their  
12 direct cytotoxicity was not the primary goal during the  
13 production of hybridomas. Any tendency for these  
14 antibodies to produce tumor cell killing is thus through  
15 chance, not by design.

16 Although the production of monoclonal antibodies have  
17 been carried out using whole cell immunization for various  
18 applications the screening of these hybridomas have relied  
19 on either putative or identified target antigens or on the  
20 selectivity of these hybridomas for specific tissues. It  
21 is axiomatic that the best antibodies are the ones with  
22 the highest binding constants. This concept originated  
23 from the basic biochemical principle that enzymes with the  
24 highest binding constants were the ones that were the most  
25 effective for catalyzing a reaction. This concept is  
26 applicable to receptor ligand binding where the drug  
27 molecule binding to the receptor with the greatest  
28 affinity usually has the highest probability for  
29 initiating or inhibiting a signal. However, this may not  
30 always be the case since it is possible that in certain  
31 situations there may be cases where the initiation or  
32 inhibition of a signal may be mediated through non-  
33 receptor binding. The information conveyed by a  
34 conformational change induced by ligand binding can have  
35 many consequences such as a signal transduction,  
36 endocytosis, among the others. The ability to produce a

1 conformational change in a receptor molecule may not  
2 necessarily be due to the filling of a ligand receptor  
3 pocket but may occur through the binding of another extra  
4 cellular domain or due to receptor clustering induced by a  
5 multivalent ligand.

6 The production of antibodies to produce cell killing  
7 need not be predicated upon screening of the hybridomas  
8 for the best binding antibodies. Rather, although not  
9 advocated by those who produce monoclonal antibodies, the  
10 screening of the hybridoma supernatants for cell killing  
11 or alternatively for cessation of growth of the cancerous  
12 cells may be selected as a desirable endpoint for the  
13 production of cytotoxic or cytostatic antibodies. It is  
14 well understood that the *in-vivo* antibodies mediate their  
15 function through the Fc portions and that the utility of  
16 the therapeutic antibody is determined by the  
17 functionality of the constant region or attached moieties.  
18 In this case the FAb portion of the antibody, the antigen-  
19 combining portion, will confer to the antibody its  
20 specificity and the Fc portion its functionality. The  
21 antigen combining site of the antibody can be considered  
22 to be the product of a natural combinatorial library. The  
23 result of the rearrangement of the variable region of the  
24 antibody can be considered a molecular combinatorial  
25 library where the output is a peptide. Therefore, the  
26 sampling of this combinatorial library can be based on any  
27 parameter. Like sampling a natural compound library for  
28 antibiotics, it is possible to sample an antibody library  
29 for cytotoxic or cytostatic compounds.

30 The various endpoints in a screen must be  
31 differentiated from each other. For example, the  
32 difference between antibody binding to the cell is  
33 distinct from cell killing. Cell killing (cytotoxicity) is  
34 distinct from the mechanisms of cell death such as oncosis  
35 or apoptosis. There can be many processes by which cell  
36 death is achieved and some of these can lead either to

1 oncosis or apoptosis. There is speculation that there are  
2 other cell death mechanisms other than oncosis or  
3 apoptosis but regardless of how the cell arrives at death  
4 there are some commonalities of cell death. One of these  
5 is the absence of metabolism and another is the  
6 denaturation of enzymes. In either case vital stains will  
7 fail to stain these cells. These endpoints of cell death  
8 have been long understood and predate the current  
9 understanding of the mechanisms of cell death.  
10 Furthermore, there is the distinction between cytotoxic  
11 effects where cells are killed and cytostatic effects  
12 where the proliferation of cells are inhibited.

13 In a preferred embodiment of the present invention,  
14 the assay is conducted by focusing on cytotoxic activity  
15 toward cancerous cells as an end point. In a preferred  
16 embodiment, a live /dead assay kit , for example the  
17 LIVE/DEAD® Viability/Cytotoxicity Assay Kit (L-3224) by  
18 Molecular Probes, is utilized. The Molecular Probes kit  
19 provides a two-color fluorescence cell viability assay  
20 that is based on the simultaneous determination of live  
21 and dead cells with two probes that measure two recognized  
22 parameters of cell viability - intracellular esterase  
23 activity and plasma membrane integrity. The assay  
24 principles are general and applicable to most eukaryotic  
25 cell types, including adherent cells and certain tissues,  
26 but not to bacteria or yeast. This fluorescence-based  
27 method of assessing cell viability is preferred in place  
28 of such assays as trypan blue exclusion, Cr release and  
29 similar methods for determining cell viability and  
30 cytotoxicity.

31 In carrying out the assay, live cells are  
32 distinguished by the presence of ubiquitous intracellular  
33 esterase activity, determined by the enzymatic conversion  
34 of the virtually nonfluorescent cell-permeant CALCEIN AM  
35 to the intensely fluorescent Calcein. The polyanionic dye  
36 Calcein is well retained within live cells, producing an



1 intense uniform green fluorescence in live cells (ex/em  
2 ~495 nm/~515 nm). EthD-1 enters cells with damaged  
3 membranes and undergoes a 40-fold enhancement of  
4 fluorescence upon binding to nucleic acids, thereby  
5 producing a bright red fluorescence in dead cells (ex/em  
6 ~495 nm/~635 nm). EthD-1 is excluded by the intact plasma  
7 membrane of live cells. The determination of cell  
8 viability depends on these physical and biochemical  
9 properties of cells. Cytotoxic events that do not affect  
10 these cell properties may not be accurately assessed using  
11 this method. Background fluorescence levels are inherently  
12 low with this assay technique because the dyes are  
13 virtually nonfluorescent before interacting with cells.

14 In addition to the various endpoints for screening,  
15 there are two other major characteristics of the screening  
16 process. The library of antibody gene products is not a  
17 random library but is the product of a biasing procedure.  
18 In the examples below, the biasing is produced by  
19 immunizing mice with fixed cells. This increases the  
20 proportion of antibodies that have the potential to bind  
21 the target antigen. Although immunization is thought of as  
22 a way to produce higher affinity antibodies (affinity  
23 maturation) in this case it is not. Rather, it can be  
24 considered as a way to shift the set of antigen combining  
25 sites towards the targets. This is also distinct from the  
26 concept of isotype switching where the functionality, as  
27 dictated by the constant portion of the heavy chain, is  
28 altered from the initial IgM isotype to another isotype  
29 such as IgG.

30 The third key feature that is crucial in the  
31 screening process is the use of multitarget screening. To  
32 a certain extent specificity is related to affinity. An  
33 example of this is the situation where an antigen has very  
34 limited tissue distribution and the affinity of the  
35 antibody is a key determinant of the specificity of the

1 antibody-the higher the affinity the more tissue specific  
2 the antibody and likewise an antibody with low affinity  
3 may bind to tissues other than the one of interest.  
4 Therefore, to address the specificity issue the antibodies  
5 are screened simultaneously against a variety of cells. In  
6 the examples below the hybridoma supernatants  
7 (representing the earliest stages of monoclonal antibody  
8 development), are tested against a number of cell lines to  
9 establish specificity as well as activity.

10 The antibodies are designed for therapeutic treatment  
11 of cancer in patients. Ideally the antibodies can be naked  
12 antibodies. They can also be conjugated to toxins. They  
13 can be used to target other molecules to the cancer. e.g.  
14 biotin conjugated enzymes. Radioactive compounds can also  
15 be used for conjugation.

16 The antibodies can be fragmented and rearranged  
17 molecularly. For example Fv fragments can be made; sFv-  
18 single chain Fv fragments; diabodies etc.

19 It is envisioned that these antibodies can be used  
20 for diagnosis, prognosis, and monitoring of cancer. For  
21 example the patients can have blood samples drawn for shed  
22 tumor antigens which can be detected by these antibodies  
23 in different formats such as ELISA assays, rapid test  
24 panel formats etc. The antibodies can be used to stain  
25 tumor biopsies for the purposes of diagnosis. In addition  
26 a panel of therapeutic antibodies can be used to test  
27 patient samples to determine if there are any suitable  
28 antibodies for therapeutic use.

#### 29 Example one

30 In order to produce monoclonal antibodies specific  
31 for a tumor sample the method of selection of the  
32 appropriate hybridoma wells is complicated by the  
33 probability of selecting wells which will produce false  
34 positive signals. That is to say that there is the

1     likelihood of producing antibodies that can react against  
2     normal cells as well as cancer cells. To obviate this  
3     possibility one strategy is to mask the anti-normal  
4     antigen antibodies from the selection process. This can  
5     be accomplished by removing the anti-normal antibodies at  
6     the first stage of screening thereby revealing the  
7     presence of the desired antibodies. Subsequent limiting  
8     dilution cloning can delineate the clones that will not  
9     produce killing of control cells but will produce target  
10    cancer cell killing.

11        Biopsy specimens of breast, melanoma, and lung tumors  
12    were obtained and stored at  $-70^{\circ}\text{C}$  until used. Single cell  
13    suspensions were prepared and fixed with  $-30^{\circ}\text{C}$ , 70%  
14    ethanol, washed with PBS and reconstituted to an  
15    appropriate volume for injection. Balb/c mice were  
16    immunized with  $2.5 \times 10^5$ - $1 \times 10^6$  cells and boosted every third  
17    week until a final pre-fusion boost was performed three  
18    days prior to the splenectomy. The hybridomas were  
19    prepared by fusing the isolated splenocytes with Sp2/0 and  
20    NS1 myeloma partners. The supernatants from the fusions  
21    were tested for subcloning of the hybridomas.

22    Cells (including A2058 melanoma cells, CCD-12CoN  
23    fibroblasts, MCF-12A breast cells among others) were  
24    obtained from ATCC and cultured according to enclosed  
25    instructions. The HEY cell line was a gift from Dr. Inka  
26    Brockhausen. The non-cancer cells, e.g. CCD-12CoN  
27    fibroblasts and MCF-12A breast cells, were plated into 96-  
28    well microtitre plates (NUNC) 1 to 2 weeks prior to  
29    screening. The cancer cells, e.g. HEY, A2058, BT 483, and  
30    HS294t, were plated two or three days prior to screening.

31        The plated normal cells were fixed prior to use. The  
32    plates were washed with 100 microliters of PBS for 10  
33    minutes at room temperature and then aspirated dry. 75  
34    microliters of 0.01 percent glutaraldehyde diluted in PBS  
35    were added to each well for five minutes and then  
36    aspirated. The plates were washed with 100 microliters of

1     PBS three times at room temperature. The wells were  
2     emptied and 100 microliters of one percent human serum  
3     albumin in PBS was added to each well for one hour at room  
4     temperature. The plates were then stored at four degrees  
5     Celsius.

6             Prior to the transfer of the supernatant from the  
7     hybridoma plates the fixed normal cells were washed three  
8     times with 100 microliters of PBS at room temperature.  
9     After aspiration to the microliters of the primary  
10    hybridoma culture supernatants were transferred to the  
11    fixed cell plates and incubated for two hours at 37  
12    degrees Celsius in a 8 percent CO<sub>2</sub> incubator. The  
13    hybridoma supernatants derived from melanoma was incubated  
14    with CCD-12 CoN cells and those derived from breast cancer  
15    were incubated with MCF-12a cells. After incubation  
16    the absorbed supernatant was divided into two 75  
17    microliter portions and transferred to target cancer cell  
18    plates. Prior to the transfer the cancer cell plates were  
19    washed three times with 100 microliters of PBS. The  
20    supernatant from the CCD-12 CoN cells were transferred to  
21    the A2058 and the HS294t cells, whereas the supernatant  
22    from MCF-12A cells were transferred to the HEY and BT 483  
23    cells. The cancer cells were incubated with the hybridoma  
24    supernatants for 18 hours at 37 degrees Celsius in an 8  
25    percent CO<sub>2</sub> incubator.

26            The Live/Dead cytotoxicity assay was obtained from  
27    Molecular Probes (Eu,OR). The assays were performed  
28    according to the manufacturer's instructions with the  
29    changes outlined below. The plates with the cells were  
30    washed once with 100 microliters of PBS at 37°C. 75 to 100  
31    microliters of supernatant from the hybridoma microtitre  
32    plates were transferred to the cell plates and incubated  
33    in a 8% CO<sub>2</sub> incubator for 18-24 hours. Then, the wells that  
34    served as the all dead control were aspirated until empty  
35    and 50 microliters of 70% ethanol was added. The plate was  
36    then emptied by inverting and blotted dry. Room

1 temperature PBS was dispensed into each well from a  
2 multichannel squeeze bottle, tapped three times, emptied  
3 by inversion and then blotted dry. 50 microliters of the  
4 fluorescent Live/Dead dye diluted in PBS was added to each  
5 well and incubated at 37°C in a 5% CO<sub>2</sub> incubator for one  
6 hour. The plates were read in a Perkin-Elmer HTS7000  
7 fluorescence plate reader and the data was analyzed in  
8 Microsoft Excel.

9 Four rounds of screening were conducted to produce  
10 single clone hybridoma cultures. For two rounds of  
11 screening the hybridoma supernatants were tested only  
12 against the cancer cells. In the last round of screening  
13 the supernatant was tested against a number of non-cancer  
14 cells as well as the target cells indicated in table 1.  
15 The antibodies were isotyped using a commercial isotyping  
16 kit.

17 A number of monoclonal antibodies were produced in  
18 accordance with the method of the present invention.  
19 These antibodies, whose characteristics are summarized in  
20 Table 1, are identified as 3BD-3, 3BD-6, 3BD-8, 3BD-9,  
21 3BD-15, 3BD-25, 3BD-26 and 3BD-27. Each of the designated  
22 antibodies is produced by a hybridoma cell line deposited  
23 with the American Type Culture Collection at 10801  
24 University Boulevard, Manassas, Va. having an ATCC  
25 Accession Number as follows:

26	<u>Antibody</u>	<u>ATCC Accession Number</u>
27	3BD-3	
28	3BD-6	
29	3BD-8	
30	3BD-9	
31	3BD-15	
32	3BD-25	
33	3BD-26	
34	3BD-27	

1 These antibodies are considered monoclonal after four  
 2 rounds of limiting dilution cloning. The anti-melanoma  
 3 antibodies did not produce significant cancer cell  
 4 killing. The panel of anti-breast cancer antibodies killed  
 5 32-87% of the target cells and <1-3% of the control cells.  
 6 The predominant isotype was IgG1 even though it was  
 7 expected that the majority of anti-tumor antibodies would  
 8 be directed against carbohydrate antigens, and thus, be of  
 9 the IgM type. There is a high therapeutic index since most  
 10 antibodies spare the control cells from cell death.

11 Table 1. Anti-Breast Cancer Antibodies

12

Clones	% Cell Death			
	Target for Anti-Breast Cancer Antibodies (HEY & A2058)	Normal Fibroblast Cells (CCD-12CoN)	Fibrocystic Breast Cells (MCF-12A)	Isotype
15 3BD-3	74.9%	3.7%	<1%	$\gamma 1, \lambda$
16 3BD-6	68.5%	5.6%	<1%	$\gamma 1, \lambda$
17 3BD-8	81.9%	4.5%	2.6%	$\gamma 1, \kappa$
18 3BD-9	77.2%	7.9%	<1%	$\gamma 1, \lambda$
19 3BD-15	87.1%	<1 %	<1%	$\gamma 1, \lambda$
20 3BD-26	54.8%	3.3%	<1%	$\mu, \kappa$
21 3BD-25	32.4%	3.6%	<1 %	$\gamma 1, \kappa$
22 3BD-27	60.1%	8.3%	1.3%	$\gamma 1, \kappa$

23

24 **Example 2**

25 In this example customized anti-cancer antibodies are  
 26 produced by first obtaining samples of the patient's  
 27 tumor. Usually this is from a biopsy specimen from a  
 28 solid tumor or a blood sample from hematogenous tumors.  
 29 The samples are prepared into single cell suspensions and  
 30 fixed for injection into mice. After the completion of the  
 31 immunization schedule the hybridomas are produced from the  
 32 splenocytes. The hybridomas are screened against a variety

1 of cancer cell lines and normal cells in standard  
2 cytotoxicity assays. Those hybridomas that are reactive  
3 against cancer cell lines but are not reactive against  
4 normal non-transformed cells are selected for further  
5 propagation. Clones that were considered positive were  
6 ones that selectively killed the cancer cells but did not  
7 kill the non-transformed cells. The antibodies are  
8 characterized for a large number of biochemical parameters  
9 and then humanized for therapeutic use.

10 The melanoma tumor cells isolated and cell lines were  
11 cultured as described in Example 1. Balb/c mice were  
12 immunized according to the following schedule: 200,000  
13 cells s.c. and i.p. on day 0, then 200,000 cells were  
14 injected i.p. on day 21, then 1,000,000 cells were  
15 injected on day 49, then 1,250,000 cells in Freund's  
16 Complete Adjuvant were injected i.p. on day 107, and then  
17 200,000 cells were injected on day 120 i.p. and then the  
18 mice were sacrificed on day 123. The spleens were  
19 harvested and the splenocytes were divided into two  
20 aliquots for fusion with Sp2/0 (1LN) or NS-1 (2LN) myeloma  
21 partners using the methods outlined in example 1.

22 The screening was carried out 11 days after the  
23 fusion against A2058 melanoma cells and CCD-12CoN  
24 fibroblasts. Each pair of plates were washed with 100  
25 microliters of room temperature PBS and then aspirated to  
26 near dryness. Then 50 microliters of hybridoma supernatant  
27 was added to the same wells on each of the two plates. The  
28 spent Sp2/0 supernatant was added to the control wells at  
29 the same volume and the plates were incubated for around  
30 18 hours at 37 degrees Celsius at a 8%CO<sub>2</sub>, 98% relative  
31 humidity incubator. Then each pair of plates were removed  
32 and in the positive control wells 50 microliters of 70%  
33 ethanol was substituted for the media for 4 seconds. The  
34 plates were then inverted and washed with room temperature  
35 PBS once and dried. Then 50uL of fluorescent live/dead dye  
36 diluted in PBS (Molecular Probes Live/Dead Kit) was added

1 for one hour and incubated at 37 degrees Celsius. The  
2 plates were then read in a Perkin-Elmer fluorescent plate  
3 reader and the data analyzed using Microsoft Excel. The  
4 wells that were considered positive were subcloned and the  
5 same screening process was repeated 13 days later and then  
6 33 days later. The results of the last screening is  
7 outlined in Table 2 below. A number of monoclonal  
8 antibodies were produced in accordance with the method of  
9 the present invention. These antibodies, whose  
10 characteristics are summarized in Table 2, are identified  
11 as 1LN-1, 1LN-8, 1LN-12, 1LN-14, 2LN-21, 2LN-28, 2LN-29,  
12 2LN-31, 2LN-33, 2LN-34 and 2LN-35. Each of the designated  
13 antibodies is produced by a hybridoma cell line deposited  
14 with the American Type Culture Collection at 10801  
15 University Boulevard, Manassas, Va. having an ATCC  
16 Accession Number as follows:

17		
18		
19	<u>Antibody</u>	<u>ATCC Accession Number</u>
20	1LN-1	
21	1LN-8	
22	1LN-12	
23	1LN-14	
24	2LN-21	
25	2LN-28	
26	2LN-29	
27	2LN-31	
28	2LN-33	
29	2LN-34	
30	2LN-35	



Table 2, Anti-Melanoma Antibodies

Clones	% Cell Death	
	Target for Anti-Melanoma Antibodies (A2058)	Normal Fibroblast Cells (CCD-1 2CoN)
1LN-1	59.4%	<1 %
1LN-8	11.0%	5.0%
1LN-12	55.2%	1.4%
1LN-14	51.4%	<1%
2LN-21	72.0%	15.9%
2LN-28	66.6%	12.4%
2LN-29	78.2%	6.1%
2LN-31	100%	7.8%
2LN-33	94.2%	<1%
2LN-34	56.6%	11.2%
2LN-35	66.5%	6.6%

The table illustrates that clones from both the Sp2/0 and NS-1 fusions were able to produce antibodies that had a greater than 50% killing rate for cancerous cells and at the same time some of the clones were able to produce less than one percent killing of normal control fibroblasts.

### Example 3

In this example antibodies were produced to several different breast tumor samples following the method of Example 2 in order to demonstrate the generality of producing customized antibodies. Biopsy specimens of breast tumors were obtained and stored at -70°C until used as noted in Example 1. Single cell suspensions were prepared for each specimen and fixed with -30°C, 70% ethanol, washed with PBS and reconstituted to an appropriate volume for injection. Female, 7-8 week old, A strain, H-2<sup>d</sup> haplotype Balb/c mice (Charles River Canada,

1 St. Constant, QC, Can), were immunized with  $2.5 \times 10^5$ - $1 \times 10^6$   
2 cells and boosted every third week until a final pre-  
3 fusion boost was performed three days prior to the  
4 splenectomy. The hybridomas were prepared by fusing the  
5 isolated splenocytes with Sp2/0 myeloma partners. The  
6 supernatants from the fusions were tested for subcloning  
7 of the hybridomas.

8  
9 Hs574.T breast ductal carcinoma cells, A2058  
10 melanoma cells, NCI-H460 human lung large cell carcinoma,  
11 NCI-H661 human lung large cell carcinoma, CCD-112CoN human  
12 colon fibroblasts, CCD-27sk human skin fibroblasts, MCF-  
13 12A human mammary epithelial cells, Hs574.mg human breast  
14 cells and other cell lines, were obtained from ATCC and  
15 cultured according to enclosed instructions. Both cancer  
16 and non-cancer cells were plated three to four days prior  
17 to screening.

18 The hybridomas were cultured for ten to twelve days  
19 after fusion and observed under the microscope. When 20 to  
20 25% of the wells were over 80% confluent, the hybridoma  
21 supernatants were screened in a cytotoxicity assay. The  
22 hybridoma supernatants were divided into two 75 microliter  
23 portions; one portion was added to a target cancer cell  
24 plate and another to a non-cancer cell plate. Prior to  
25 transfer of hybridoma supernatants, the cell plates were  
26 washed three times with 100 microliters of PBS. The  
27 supernatant from the anti-breast cancer hybridomas were  
28 transferred to the Hs574.T and the Hs574.mg cells, whereas  
29 the supernatant from the anti-lung cancer hybridoma were  
30 transferred to the NCI-H460 and CCD-27SK cells. The

1 cancer cells were incubated with the hybridoma  
2 supernatants for 18 hours at 37 degrees Celsius in an 8  
3 percent CO<sub>2</sub> incubator.

4 The Live/Dead cytotoxicity assay was obtained from  
5 Molecular Probes (Eugene, OR). The assays were performed  
6 according to the manufacturer's instructions with the  
7 changes outlined below. The plates with the cells were  
8 washed once with 100 microliters of PBS at 37°C. 75 to 100  
9 microliters of supernatant from the hybridoma microtitre  
10 plates were transferred to the cell plates and incubated  
11 in a 8% CO<sub>2</sub> incubator for 18-24 hours. Then, the wells that  
12 served as the dead control cells were aspirated until  
13 empty and 50 microliters of 70% ethanol was added. The  
14 plate was then emptied by inverting and blotted dry. Room  
15 temperature PBS was dispensed into each well from a  
16 multichannel squeeze bottle, tapped three times, emptied  
17 by inversion and then blotted dry. 50 microliters of the  
18 fluorescent Live/Dead dye diluted in PBS was added to each  
19 well and incubated at 37°C in a 5% CO<sub>2</sub> incubator for one  
20 hour. The plates were read in a Perkin-Elmer HTS7000  
21 fluorescence plate reader and the data was analyzed in  
22 Microsoft Excel (Microsoft, Redmond, WA).

23 Four rounds of screening were conducted to  
24 produce single clone hybridoma cultures. For two rounds of  
25 screening the hybridoma supernatants were tested only  
26 against the cancer cells. In the last round of screening  
27 the supernatant was tested against a number of non-cancer

1 cells as well as the target cells indicated in Table 3.  
2 The antibodies were isotyped using a commercial isotyping  
3 kit (Roche, Indianapolis, IN).

4 A number of monoclonal antibodies were produced in  
5 accordance with the method of the present invention.  
6 These antibodies, whose characteristics are summarized in  
7 Table 3, are identified as 4BD-1, 4BD-3, 4BD-6, 4BD-9,  
8 4BD-13, 4BD-18, 4BD-20, 4BD-25, 4BD-37, 4BD-32, 4BD-26,  
9 4BD-27, 4BD-28, 4BD-50, 6BD-1, 6BD-3, 6BD-5, 6BD-11, 6BD-  
10 25, 7BD-7, 7BD-12-1, 7BD-12-2, 7BD-13, 7BD-14, 7BD-19,  
11 7BD-21, 7BD-24, 7BD-29, 7BD-30, 7BD-31, 7BDI-17, 7BDI-58,  
12 7BDI-60 and 7BDI-62. Each of the designated antibodies is  
13 produced by a hybridoma cell line deposited with the  
14 American Type Culture Collection at 10801 University  
15 Boulevard, Manassas, Va. having an ATCC Accession Number  
16 as follows:

17	<u>Antibody</u>	<u>ATCC Accession Number</u>
18	4BD-1	
19	4BD-3	
20	4BD-6	
21	4BD-9	
22	4BD-13	
23	4BD-18	
24	4BD-20	
25	4BD-25	
26	4BD-37	
27	4BD-32	

- 1 4BD-26
- 2 4BD-27
- 3 4BD-28
- 4 4BD-50
- 5 6BD-1
- 6 6BD-3
- 7 6BD-5
- 8 6BD-11
- 9 6BD-25
- 10 7BD-7
- 11 7BD-12-1
- 12 7BD-12-2
- 13 7BD-13
- 14 7BD-14
- 15 7BD-19
- 16 7BD-21
- 17 7BD-24
- 18 7BD-29
- 19 7BD-30
- 20 7BD-31
- 21 7BDI-17
- 22 7BDI-58
- 23 7BDI-60
- 24 7BDI-62

25        These antibodies are considered monoclonal after four  
26 rounds of limiting dilution cloning. The panel of anti-  
27 breast cancer antibodies killed 15-79% of the target cells  
28 and <1-31% of the control cells. The majority of anti-

1 tumor antibodies were IgM type, suggesting they could be  
2 directed against carbohydrate antigens on the surface of  
3 tumor cells. There is a high therapeutic index since most  
4 of the antibodies do not cause the normal cells to undergo  
5 cell death.

6 These monoclonal antibodies are characterized for a  
7 number of immunological and biochemical parameters. A  
8 cell based enzyme linked immunosorbent assay (ELISA) was  
9 established for measuring the binding of the antibodies  
10 derived of each clones to different cell lines. Cells were  
11 seeded and grown on 96-well tissue culture plates. The  
12 plates were washed with 100 microliters of PBS. 100  
13 microliters of cold 4 percent paraformaldehyde in PBS were  
14 added to each well for ten minutes and then aspirated. The  
15 plates were washed with PBS using a multichannel squeeze  
16 bottle . The wells were emptied and 100 microliters of  
17 blocking buffer (1 percent hydrocasein, 0.1 percent  
18 geletin in 50mM Tris-HCl buffer, pH 9.3) was added to each  
19 well for one hour at room temperature. The plates were  
20 washed three times with a buffer (0.05 percent Tween 20 in  
21 10 mM PBS) at room temperature and then stored at -30  
22 degrees Celsius with 100 microliters of the buffer. Prior  
23 to use the plates were thawed and the buffer was aspirated  
24 from each well. 75 microliters of hybridoma supernatant  
25 were added to each well and incubated for 60 minutes at  
26 room temperature. After the plates were washed with PBS  
27 using a multichannel squeeze bottle, 50 microliters of a

1 combination of peroxidase conjugated goat anti-mouse IgG  
2 and peroxidase conjugated donkey anti-mouse IgM (Jackson  
3 ImmunoResearch Lab, Inc., West Grove, PA.) was added and  
4 incubated for 30 minutes at room temperature. After the  
5 last wash, 50 microliters of orthophenylene diamine (OPD)  
6 (Sigma, St. Louis, MO) was added to each well and the  
7 optical density was read at 492 nm on the HTS7000 plate  
8 reader after adding equal volume of 1 N sulfuric acid.  
9 Different clones show different profiles in binding to  
10 different cells (Table 3). This indicates that they may  
11 target different cell surface antigen and further suggests  
12 the variable distribution of these antigen on the surface  
13 of cancer cells. Those binding only to cancer cells but  
14 not to normal cells could identify certain tumor-  
15 associated antigen.

16

17 Table 3. Anti-Breast Cancer Antibodies

Clones	Isotype	% Cell Death		Binding to cell lines				
		Hs574.T	Hs574.mg	Hs574.T	Hs574.mg	NCI-H460	CCD-27sk	A2058
6BD-1	$\mu, \kappa$	38.2	5	0.8	0.5	0.6	0.3	ND*
6BD-3	$\mu, \kappa$	79	12	0.35	0.25	0.24	0.14	ND
6BD-5	$\mu, \kappa$	57.3	8	1.0	0.3	0.14	0.25	ND
6BD-11	$\mu, \kappa$	52.3	11	0.15	0.1	0.17	0.1	ND
6BD-25	$\mu, \kappa$	33.3	2	0.15	0.1	0.2	0.1	ND
4BD-26	$\mu, \kappa$	27	1.8	0.5	ND	ND	<0.1	ND
4BD-27	$\mu, \kappa$	19.6	<1	0.9	ND	ND	0.5	ND
4BD-28	$\mu, \kappa$	26.4	<1	0.8	ND	ND	<0.1	ND
4BD-32	$\mu, \kappa$	41.7	4	0.8	ND	ND	<0.1	ND
4BD-50	$\mu, \kappa$	20	<1	0.8	ND	ND	<0.1	ND
4BD-1	$\mu, \kappa$	23	31	0.6	ND	ND	<0.1	ND
4BD-3	$\mu, \kappa$	29.7	8.2	1.7	ND	ND	0.1	ND
4BD-6	$\mu, \kappa$	17	<1	0.9	ND	ND	<0.1	ND
4BD-9	$\mu, \kappa$	15	<1	0.6	ND	ND	<0.1	ND
4BD-13	$\mu, \kappa$	31	<1	1.2	ND	ND	<0.1	ND
4BD-18	$\mu, \kappa$	23.3	2.4	0.7	ND	ND	0.12	ND
4BD-20	$\mu, \kappa$	45	<1	0.95	ND	ND	<0.1	ND
4BD-25	$\mu, \kappa$	26	14.16	1.8	ND	ND	0.1	ND
4BD-37	$\mu, \kappa$	30	<1	0.8	ND	ND	<0.1	ND
7BD-7	$\mu, \kappa$	24	3	0.8	0.3	1.4	0.26	ND
7BD-12-1	$\mu, \kappa$	22	6	0.36	0.16	0.43	0.1	ND
7BD-12-2	$\mu, \kappa$	31	2	0.2	0.2	0.2	0.2	0.2
7BD-13	$\mu, \kappa$	29	12	0.1	0.15	0.2	0.1	0.2
7BD-14	$\mu, \kappa$	32	13	0.4	0.4	0.6	0.3	0.5
7BD-19	$\mu, \kappa$	20	4	1.3	0.4	0.43	0.2	ND
7BD-21	$\mu, \kappa$	21	13	0.4	0.5	0.25	0.3	ND
7BD-24	$\mu, \kappa$	32	15	0.3	0.1	0.14	0.15	ND
7BD-29	$\mu, \kappa$	15	16	0.3	0.24	0.14	0.16	ND
7BD-30	$\mu, \kappa$	23	13	0.34	0.24	0.2	0.16	ND
7BD-31	$\mu, \kappa$	28	10	0.3	0.4	0.4	0.3	0.4



1	7BDI-17	$\mu, \kappa$	23	<1	0.75	ND	ND	ND	ND
2	7BDI-58	$\gamma 1, \kappa$	17.5	<1	0.77	ND	ND	ND	ND
3	7BDI-60	$\gamma 1, \kappa$	15	<1	0.73	ND	ND	ND	ND
4	7BDI-62		15	5	0.55	ND	ND	ND	ND

5 \*ND: not done.

#### 6 Example 4

7 In this example customized anti-cancer antibodies are  
8 produced to a lung cancer sample by first obtaining  
9 samples of the patient's tumor preparing single cell  
10 suspensions which are then fixed for injection into mice  
11 as noted in Example 1. After the completion of the  
12 immunization schedule the hybridomas are produced from the  
13 splenocytes. The hybridomas are screened against a variety  
14 of cancer cell lines and normal cells in standard  
15 cytotoxicity assays. Those hybridomas that are reactive  
16 against cancer cell lines but are not reactive against  
17 normal non-transformed cells are selected for further  
18 propagation. Clones that were considered positive were  
19 ones that selectively killed the cancer cells but did not  
20 kill the non-transformed cells.

21 The lung cancer cells were isolated and cell lines  
22 were cultured as described in Example 1. Female, 7-8 week  
23 old, A strain, H-2<sup>d</sup> haplotype Balb/c mice (Charles River  
24 Canada, St. Constant, QC, Can), were immunized with human  
25 lung cancer cells. The lung cancer cell suspensions were  
26 emulsified in an equal volume of Freund's complete  
27 adjuvant (FCA) for the first immunization and then in

1 Freund's incomplete adjuvant (FIA) for subsequent  
2 immunizations at 0, 21, 45 days.  $5 \times 10^5$  cells were used to  
3 immunize each mouse either through a subcutaneous or  
4 intra-peritoneal route. Immunized mice were sacrificed 3-4  
5 days after the final immunization with human lung cancer  
6 cells at 148 days, given intra-peritoneally, in PBS at pH  
7 7.4. The spleens were harvested and the splenocytes were  
8 divided into two aliquots for fusion with Sp2/0 myeloma  
9 partners using the methods outlined in Example 1.

10 The screening was carried out 10 days after the  
11 fusion against NCI-H460 and/or NCI-H661 cells and CCD-27SK  
12 fibroblasts. Each pair of plates were washed with 100  
13 microliters of room temperature PBS and then aspirated to  
14 near dryness. Then 75 microliters of hybridoma supernatant  
15 was added per well on each of the two plates. The spent  
16 Sp2/0 supernatant was added to the control wells at the  
17 same volume and the plates were incubated for around 18  
18 hours at 37 degrees Celsius at a 8%CO<sub>2</sub>, 98% relative  
19 humidity incubator. Then each pair of plates was removed  
20 and in the positive control wells 50 microliters of 70%  
21 ethanol was substituted for the media for 4 seconds. The  
22 plates were then inverted and washed with room temperature  
23 PBS once and dried. Then 50 microliters of fluorescent  
24 live/dead dye diluted in PBS (Molecular Probes Live/Dead  
25 Kit) was added for one hour and incubated at 37 degrees  
26 Celsius. The plates were then read in a Perkin-Elmer  
27 fluorescent plate reader and the data analyzed using

1 Microsoft Excel. The wells that were considered positive  
2 were subcloned and the same screening process was repeated  
3 6 days later and then 13 days later. The result of the  
4 last screening is outlined in Table 4 below. Antibodies  
5 were characterized for binding to different cell lines  
6 with a cellular ELISA according to the methods of Example  
7 3. A number of monoclonal antibodies were produced in  
8 accordance with the method of the present invention.  
9 These antibodies, whose characteristics are summarized in  
10 Table 4, are identified as 5LAC2, 5LAC4, 5LAC20, and  
11 5LAC23. Each of the designated antibodies is produced by  
12 a hybridoma cell line deposited with the American Type  
13 Culture Collection at 10801 University Boulevard,  
14 Manassas, Va. having an ATCC Accession Number as follows:

15	<u>Antibody</u>	ATCC Accession Number

16 5LAC2

17 5LAC4

18 5LAC20

19 5LAC23.

20      Table 4. Anti-Lung Cancer Antibodies

Clones	Isotype	% Cell Death					Binding to cell lines				
		Hs574.T	NCI-H460	NCI-H661	A2058	CCD-27sk	Hs574.T	Hs574.mg	NCI-H460	CCD-27sk	A2058
5LAC2	μ, κ	30	7	45.3	23	<1	0.2	0.2	0.26	0.2	0.2
5LAC4	μ, κ	21	11	20.5	23	3	0.7	0.9	1.7	0.8	0.9
5LAC20	μ, κ	23	7	66.4	24	3	0.5	0.2	0.6	0.2	0.2
		23	8	57.6	25	5	0.6	0.6	0.6	0.6	0.6

1     \*ND: not done

2           The table illustrates that clones were able to  
3     produce antibodies that had a greater than 7-67% killing  
4     rate for cancerous cells and at the same time some of the  
5     clones were able to produce less than five percent killing  
6     of normal control fibroblasts.

7

8     **Example 5**

9           In this example customized anti-cancer antibodies are  
10    produced to a patient's lung cancer cells, but cultured  
11    cells were used in the antibody development process to  
12    demonstrate the generality of the immunization process.  
13    The samples were prepared into single cell suspensions and  
14    fixed for injection into mice as noted in Example 1. After  
15    the completion of three rounds of immunization with cells  
16    derived directly from a patient's lung cancer, the mice  
17    were immunized twice with a human lung large cell  
18    carcinoma cell line (NCI-H460). Hybridomas were produced  
19    from splenocytes and the supernatants were screened  
20    against a variety of cancer cell lines and normal cells in  
21    standard cytotoxicity assays. Those hybridomas that were  
22    reactive against cancer cell lines but were not reactive  
23    against normal non-transformed cells were selected for  
24    further propagation. Clones that were considered positive  
25    were ones that selectively killed the cancer cells but did  
26    not kill the non-transformed cells. The antibodies are

1 characterized for a large number of biochemical parameters  
2 and then humanized for therapeutic use.

3         The lung tumor cells isolated and cell lines were  
4 cultured as described in Example 1. Balb/c mice, A strain  
5 with H-2<sup>d</sup> haplotype from Charles River Canada, St.  
6 Constant, Quebec, Canada, female, 7-8 week old, were  
7 immunized with the human lung cancer cells emulsified in  
8 an equal volume of either Freund's complete adjuvant (FCA)  
9 for the first immunization and then in Freund's incomplete  
10 adjuvant (FIA) for subsequent immunizations at 0, 21, 45  
11 days with  $5 \times 10^5$  cells. The mice were immunized with fixed  
12 NCI H460 cells, which were prepared from NCI H460 cells  
13 grown in T-75 cell culture flask by scraping mono-layer  
14 cells into cell suspensions at 105, 150 and 170 days.  
15 Immunized mice were sacrificed 3-4 days after the final  
16 immunization with NCI H460 cells, given intra-  
17 peritoneally, in phosphate buffered saline buffer (PBS),  
18 pH 7.4. The spleens were harvested and the splenocytes  
19 were divided into two aliquots for fusion with Sp2/0  
20 myeloma partners using the methods outlined in Example 1.

21         The screening was carried out 10 days after the  
22 fusion against NCI H460 cells and CCD-27SK fibroblasts as  
23 described in Example 4. Antibodies were characterized for  
24 binding to different cell lines with a cellular ELISA  
25 according to the methods of Example 3.

1           The wells that were considered positive were  
2   subcloned and the same screening process was repeated 9  
3   days and 18 days later. The results are outlined in Table  
4   5 below. A number of monoclonal antibodies were produced  
5   in accordance with the method of the present invention.  
6   These antibodies, whose characteristics are summarized in  
7   Table 5, are identified as H460-1, H460-4, H460-5, H460-  
8   10, H460-14, H460-16-1, H460-16-2, H460-23 and H460-27.  
9   Each of the designated antibodies is produced by a  
10   hybridoma cell line deposited with the American Type  
11   Culture Collection at 10801 University Boulevard,  
12   Manassas, Va. having an ATCC Accession Number as follows:

13	<u>Antibody</u>	<u>ATCC Accession Number</u>
14	H460-1	
15	H460-4	
16	H460-5	
17	H460-10	
18	H460-14	
19	H460-16-1	
20	H460-16-2	
21	H460-23	
22	H460-27	
23		

Table 5. Anti-Lung Cancer Antibodies

Clones	Isotype	% Cell Death				Binding to cell lines				
		NCI-H460	Hs574.	A2058	CCD-	Hs574.	Hs574.	NCI-	CCD-	A2058
H460-1	$\gamma 1, \hat{e}$	16	30	23	<1	1.0	0.6	0.5	0.7	ND
H460-4		37	21	23	3	1.0	0.6	0.4	0.6	ND
H460-5	$\mu, \kappa$	22.5	23	24	3	1.0	0.3	0.3	0.2	ND
H460-10	$\mu, \kappa$	8	23	25	5	0.3	0.14	0.2	0.1	ND
H460-14	$\gamma 1, \hat{e}$	17	ND	ND	4	1.1	0.6	0.4	0.54	ND
H460-16-1	$\gamma 1, \hat{e}$	33	ND	ND	8	1.0	0.6	0.3	0.5	ND
H460-16-2	$\gamma 1, \hat{e}$	22	ND	ND	3	1.0	0.6	0.3	0.7	ND
H460-22-1	$\gamma 1, \hat{e}$	21	ND	ND	5	0.6	0.4	0.3	0.4	ND
H460-22-2	$\mu, \kappa$	23	ND	ND	3	0.4	0.1	0.1	0.1	ND
H460-23	$\mu, \kappa$	36	36	18	1	0.4	1.1	0.54	0.53	0.58
H460-27	$\mu, \kappa$	33	31	16	8	0.3	0.4	0.4	0.3	0.4

\*ND: not done

The table illustrates that clones were able to produce antibodies that had a greater than 15% killing rate for cancerous cells and at the same time some of the clones were able to produce less than eight percent killing of normal control fibroblasts.

The anti-cancer antibodies of the invention are useful for treating a patient with a cancerous disease when administered in admixture with a pharmaceutically acceptable adjuvant, for example normal saline, a lipid emulsion, albumen, phosphate buffered saline or the like and are administered in an amount effective to mediate treatment of said cancerous disease, for example with a

1 range of about 1 microgram per milliliter to about 1 gram  
2 per milliliter.

3 The method for treating a patient suffering from a  
4 cancerous disease may further include the use of  
5 conjugated anti-cancer antibodies and would this include  
6 conjugating patient specific anti-cancer antibodies with a  
7 member selected from the group consisting of toxins,  
8 enzymes, radioactive compounds, and hematogenous cells;  
9 and administering these conjugated antibodies to the  
10 patient; wherein said anti-cancer antibodies are  
11 administered in admixture with a pharmaceutically  
12 acceptable adjuvant, for example normal saline, a lipid  
13 emulsion, albumen, phosphate buffered saline or the like  
14 and are administered in an amount effective to mediate  
15 treatment of said cancerous disease, for example with a  
16 range of about 1 microgram per mil to about 1 gram per  
17 mil. In a particular embodiment, the anti-cancer  
18 antibodies useful in either of the above outlined methods  
19 may be a humanized antibody.

20 The anti-cancer antibodies of the invention are  
21 useful for treating a patient with a cancerous disease  
22 when administered in admixture with a pharmaceutically  
23 acceptable adjuvant, for example normal saline, a lipid  
24 emulsion, albumen, phosphate buffered saline or the like  
25 and are administered in an amount effective to mediate  
26 treatment of said cancerous disease, for example with a



1 range of about 1 microgram per mil to about 1 gram per  
2 mil.

3       The method for treating a patient suffering from a  
4 cancerous disease may further include the use of  
5 conjugated anti-cancer antibodies and would this include  
6 conjugating patient specific anti-cancer antibodies with a  
7 member selected from the group consisting of toxins,  
8 enzymes, radioactive compounds, and hematogenous cells;  
9 and  
10 administering these conjugated antibodies to the patient;  
11 wherein said anti-cancer antibodies are administered in  
12 admixture with a pharmaceutically acceptable adjuvant, for  
13 example normal saline, a lipid emulsion, albumen,  
14 phosphate buffered saline or the like and are administered  
15 in an amount effective to mediate treatment of said  
16 cancerous disease, for example with a range of about 1  
17 microgram per mil to about 1 gram per mil. In a  
18 particular embodiment, the anti-cancer antibodies useful  
19 in either of the above outlined methods may be a humanized  
20 antibody.

21

22

23

24

CLAIMS

What is claimed is:

Claim 1. A method for treating a patient suffering from a cancerous disease comprising:

administering to said patient anti-cancer antibodies or fragments thereof produced in accordance with a method for the production of individually customized anti-cancer antibodies which are useful in treating a cancerous disease, said antibodies including a subset of antibodies or fragments thereof characterized as being cytotoxic against cells of a cancerous tissue, said subset being essentially benign to non-cancerous cells;

wherein one or more antibodies or fragments thereof selected from said subset are placed in admixture with a pharmaceutically acceptable adjuvant and are administered in an amount effective to mediate treatment of said cancerous disease;

said one or more antibodies or fragments thereof being selected from the group consisting of a 1LN-8, 4BD-1, a 4BD-3, a 4BD-6, a 4BD-9, a 4BD-13, a 4BD-18, a 4BD-20, a 4BD-25, a 4BD-26, a 4BD-27, a 4BD-28, a 4BD-32, a 4BD-37, a 4BD-50, a 6BD-1, a 6BD-3, a 6BD-5, a 6BD-11, a 6BD-25, a 7BD-7, a 7BD-12-1, a 7BD-12-2, a 7BD-13, a 7BD-14, a 7BD-19, a 7BD-21, a 7BD-24, a 7BD-29, a 7BD-30, a 7BD-31, a 7BDI-17, a 7BDI-58, a 7BDI-60, a 7BDI-62, a 5LAC2, a 5LAC4, a 5LAC20, a 5LAC23, a H460-1, a H460-4, a

1 H460-5, a H460-10, a H460-14, a H460-16-1, a H460-16-2, a  
2 H460-23 and a H460-27 monoclonal antibody or combinations  
3 thereof.  
4

5 Claim 2. The method for treating a patient suffering  
6 from a cancerous disease in accordance with claim 1,  
7 wherein said one or more antibodies or fragments thereof  
8 selected from said subset are humanized.  
9

10 Claim 3. The method for treating a patient suffering  
11 from a cancerous disease in accordance with claim 1  
12 comprising:

13 conjugating said subset of antibodies or fragments  
14 thereof with a member selected from the group consisting  
15 of toxins, enzymes, radioactive compounds, and  
16 hematogenous cells; and

17 administering conjugated antibodies or fragments  
18 thereof to said patient;

19 wherein said conjugated antibodies are placed in  
20 admixture with a pharmaceutically acceptable adjuvant and  
21 are administered in an amount effective to mediate  
22 treatment of said cancerous disease.  
23

24 Claim 4. The method of claim 3, wherein said one or  
25 more antibodies or fragments thereof selected from said  
26 subset are humanized.

1           Claim 5. The method for treating a patient suffering  
2   from a cancerous disease in accordance with claim 1  
3   wherein:

4           the cytotoxicity of said antibodies or fragments  
5   thereof is mediated through antibody dependent cellular  
6   toxicity.

7  
8           Claim 6. The method for treating a patient suffering  
9   from a cancerous disease in accordance with claim 1  
10   wherein:

11           the cytotoxicity of said antibodies or fragments  
12   thereof is mediated through complement dependent cellular  
13   toxicity.

14  
15           Claim 7. The method for treating a patient suffering  
16   from a cancerous disease in accordance with claim 1  
17   wherein:

18           the cytotoxicity of said antibodies or fragments  
19   thereof is mediated through catalyzing of the hydrolysis  
20   of cellular chemical bonds.

21  
22           Claim 8. The method for treating a patient suffering  
23   from a cancerous disease in accordance with claim 1  
24   wherein:

25           the cytotoxicity of said antibodies or fragments  
26   thereof is mediated through producing an immune response  
27   against putative cancer antigens residing on tumor cells.

1           Claim 9. The method for treating a patient suffering  
2           from a cancerous disease in accordance with claim 1  
3           wherein:

4           the cytotoxicity of said antibodies or fragments  
5           thereof is mediated through targeting of cell membrane  
6           proteins to interfere with their function.

7

8           Claim 10. The method for treating a patient suffering  
9           from a cancerous disease in accordance with claim 1  
10          wherein:

11          the cytotoxicity of said antibodies or fragments  
12          thereof is mediated through production of a conformational  
13          change in a cellular protein effective to produce a signal  
14          to initiate cell-killing.

15

16          Claim 11. The method for treating a patient suffering  
17          from a cancerous disease in accordance with claim 1  
18          wherein:

19          said method of production utilizes a tissue sample  
20          containing cancerous and non-cancerous cells obtained from  
21          a particular individual.

22

23          Claim 12. A method for treating a patient suffering  
24          from a cancerous disease comprising:

25          administering to said patient anti-cancer antibodies  
26          or fragments thereof produced in accordance with a method  
27          for the production of individually customized anti-cancer  
28          antibodies which are useful in treating a cancerous

1 disease, said antibodies including a subset of antibodies  
2 or fragments thereof characterized as being cytotoxic  
3 against cells of a cancerous tissue, said subset being  
4 essentially benign to non-cancerous cells;

5 wherein one or more antibodies or fragments thereof  
6 selected from said subset are placed in admixture with a  
7 pharmaceutically acceptable adjuvant and are administered  
8 in an amount effective to mediate treatment of said  
9 cancerous disease;

10 said one or more antibodies or fragments thereof  
11 produced by a hybridoma cell line having an ATCC Accession  
12 Number selected from the group consisting of (to be  
13 provided before publication) or combinations thereof.  
14

15 Claim 13. The method for treating a patient suffering  
16 from a cancerous disease in accordance with claim 12,  
17 wherein said one or more antibodies or fragments thereof  
18 selected from said subset are humanized.  
19

20 Claim 14. The method for treating a patient suffering  
21 from a cancerous disease in accordance with claim 12  
22 comprising:

23 conjugating said subset of antibodies or fragments  
24 thereof with a member selected from the group consisting  
25 of toxins, enzymes, radioactive compounds, and  
26 hematogenous cells; and

27 administering conjugated antibodies or fragments  
28 thereof to said patient;

1            wherein said conjugated antibodies are placed in  
2            admixture with a pharmaceutically acceptable adjuvant and  
3            are administered in an amount effective to mediate  
4            treatment of said cancerous disease.

5  
6            Claim 15. The method of claim 14, wherein said one or  
7            more antibodies or fragments thereof selected from said  
8            subset are humanized.

9  
10           Claim 16. The method for treating a patient suffering  
11           from a cancerous disease in accordance with claim 12  
12           wherein:

13           the cytotoxicity of said antibodies or fragments  
14           thereof is mediated through antibody dependent cellular  
15           toxicity.

16  
17           Claim 17. The method for treating a patient suffering  
18           from a cancerous disease in accordance with claim 12  
19           wherein:

20           the cytotoxicity of said antibodies or fragments  
21           thereof is mediated through complement dependent cellular  
22           toxicity.

23  
24           Claim 18. The method for treating a patient suffering  
25           from a cancerous disease in accordance with claim 12  
26           wherein:

27

1           the cytotoxicity of said antibodies or fragments  
2   thereof is mediated through catalyzing of the hydrolysis  
3   of cellular chemical bonds.

4  
5           Claim 19. The method for treating a patient suffering  
6   from a cancerous disease in accordance with claim 12  
7   wherein:

8           the cytotoxicity of said antibodies or fragments  
9   thereof is mediated through producing an immune response  
10   against putative cancer antigens residing on tumor cells.

11  
12           Claim 20. The method for treating a patient suffering  
13   from a cancerous disease in accordance with claim 12  
14   wherein:

15           the cytotoxicity of said antibodies or fragments  
16   thereof is mediated through targeting of cell membrane  
17   proteins to interfere with their function.

18  
19           Claim 21. The method for treating a patient suffering  
20   from a cancerous disease in accordance with claim 12  
21   wherein:

22           the cytotoxicity of said antibodies or fragments  
23   thereof is mediated through production of a conformational  
24   change in a cellular protein effective to produce a signal  
25   to initiate cell-killing.

26  
27



1           Claim 22. The method for treating a patient suffering  
2           from a cancerous disease in accordance with claim 12  
3           wherein:

4           said method of production utilizes a tissue sample  
5           containing cancerous and non-cancerous cells obtained from  
6           a particular individual.

7  
8  
9  
10          Claim 23. Anti-cancer antibodies or fragments thereof  
11         selected from the group consisting of a 1LN-8, 4BD-1, a  
12         4BD-3, a 4BD-6, a 4BD-9, a 4BD-13, a 4BD-18, a 4BD-20, a  
13         4BD-25, a 4BD-26, a 4BD-27, a 4BD-28, a 4BD-32, a 4BD-37,  
14         a 4BD-50, a 6BD-1, a 6BD-3, a 6BD-5, a 6BD-11, a 6BD-25, a  
15         7BD-7, a 7BD-12-1, a 7BD-12-2, a 7BD-13, a 7BD-14, a 7BD-  
16         19, a 7BD-21, a 7BD-24, a 7BD-29, a 7BD-30, a 7BD-31, a  
17         7BDI-17, a 7BDI-58, a 7BDI-60, a 7BDI-62, a 5LAC2, a  
18         5LAC4, a 5LAC20, a 5LAC23, a H460-1, a H460-4, a H460-5, a  
19         H460-10, a H460-14, a H460-16-1, a H460-16-2, a H460-23  
20         and a H460-27 monoclonal antibody or combinations thereof.

21  
22  
23          Claim 24. Anti-cancer antibodies or fragments thereof  
24         produced by a hybridoma cell line having an ATCC Accession  
25         Number selected from the group  
26         consisting of           (to be provided before publication).

27  
28

1           Claim 25. The use of a composition for treating a  
2   patient suffering from a cancerous disease by  
3   administration of an effective amount of said composition  
4   to a patient to mediate treatment of said cancerous  
5   disease, wherein said composition comprises one or more  
6   antibodies or fragments thereof selected from a subset of  
7   said antibodies or fragments in admixture with a  
8   pharmaceutically acceptable adjuvant, said anti-cancer  
9   antibodies or fragments thereof produced in accordance  
10   with a method for the production of individually  
11   customized anti-cancer antibodies which are useful in  
12   treating cancerous disease, said subset of antibodies or  
13   fragments thereof characterized as being cytotoxic against  
14   cells of a cancerous tissue, as being essentially benign  
15   to non-cancerous cells and being selected from the group  
16   consisting of a 1LN-8, 4BD-1, a 4BD-3, a 4BD-6, a 4BD-9, a  
17   4BD-13, a 4BD-18, a 4BD-20, a 4BD-25, a 4BD-26, a 4BD-27,  
18   a 4BD-28, a 4BD-32, a 4BD-37, a 4BD-50, a 6BD-1, a 6BD-3,  
19   a 6BD-5, a 6BD-11, a 6BD-25, a 7BD-7, a 7BD-12-1, a 7BD-  
20   12-2, a 7BD-13, a 7BD-14, a 7BD-19, a 7BD-21, a 7BD-24, a  
21   7BD-29, a 7BD-30, a 7BD-31, a 7BDI-17, a 7BDI-58, a 7BDI-  
22   60, a 7BDI-62, a 5LAC2, a 5LAC4, a 5LAC20, a 5LAC23, a  
23   H460-1, a H460-4, a H460-5, a H460-10, a H460-14, a H460-  
24   16-1, a H460-16-2, a H460-23 and a H460-27 monoclonal  
25   antibody or combinations thereof.

26

27

1           Claim 26. The use of a composition for treating a  
2   patient suffering from a cancerous disease by  
3   administration of an effective amount of the composition  
4   to the patient to mediate treatment of said cancerous  
5   disease, wherein said composition comprises one or more  
6   antibodies or fragments thereof from a subset of  
7   antibodies or fragments thereof characterized as being  
8   cytotoxic against cells of a cancerous tissue and  
9   essentially benign to non-cancerous cells placed in  
10   admixture with a pharmaceutically acceptable adjuvant,  
11   said one or more antibodies or fragments thereof produced  
12   by a hybridoma cell line having an ATCC Accession Number  
13   selected from the group consisting of (to be provided  
14   before publication) or combinations thereof.

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original claims 1-26 replaced by amended claims 1-25 (04 pages).

1

2       What is claimed is:

3

4       Claim 1. A method for treating a patient suffering  
5 from a cancerous disease comprising:6       administering to said patient anti-cancer antibodies  
7 or fragments thereof produced in accordance with a method  
8 for the production of individually customized anti-cancer  
9 antibodies which are useful in treating a cancerous  
10 disease, said antibodies including a subset of antibodies  
11 or fragments thereof characterized as being cytotoxic  
12 against cells of a cancerous tissue, said subset being  
13 essentially benign to non-cancerous cells;14       wherein one or more antibodies or fragments thereof  
15 selected from said subset are placed in admixture with a  
16 pharmaceutically acceptable adjuvant and are administered  
17 in an amount effective to mediate treatment of said  
18 cancerous disease;19       said one or more antibodies or fragments thereof  
20 being selected from the group consisting of a 1LN-8, 4BD-  
21 1, a 4BD-3, a 4BD-6, a 4BD-9, a 4BD-13, a 4BD-18, a 4BD-  
22 20, a 4BD-25, a 4BD-26, a 4BD-27, a 4BD-28, a 4BD-32, a  
23 4BD-37, a 4BD-50, a 6BD-1, a 6BD-3, a 6BD-5, a 6BD-11, a  
24 6BD-25, a 7BD-7, a 7BD-12-1, a 7BD-12-2, a 7BD-13, a 7BD-  
25 14, a 7BD-19, a 7BD-21, a 7BD-24, a 7BD-29, a 7BD-30, a  
26 7BD-31, a 7BDI-17, a 7BDI-58, a 7BDI-60, a 7BDI-62, a  
27 5LAC2, a 5LAC4, a 5LAC20, a 5LAC23, a H460-1, a H460-4, a

1 H460-5, a H460-10, a H460-14, a H460-16-1, a H460-16-2,  
2 a H460-22-1, a H460-23 and a H460-27 monoclonal antibody or  
3 combinations thereof.  
4

5 Claim 2. The method for treating a patient suffering  
6 from a cancerous disease in accordance with claim 1,  
7 wherein said one or more antibodies or fragments thereof  
8 selected from said subset are humanized.  
9

10 Claim 3. The method for treating a patient suffering  
11 from a cancerous disease in accordance with claim 1  
12 comprising:

13 conjugating said subset of antibodies or fragments  
14 thereof with a member selected from the group consisting  
15 of toxins, enzymes, radioactive compounds, and  
16 hematogenous cells; and

17 administering conjugated antibodies or fragments  
18 thereof to said patient;

19 wherein said conjugated antibodies are placed in  
20 admixture with a pharmaceutically acceptable adjuvant and  
21 are administered in an amount effective to mediate  
22 treatment of said cancerous disease.  
23

24 Claim 4. The method of claim 3, wherein said one or  
25 more antibodies or fragments thereof selected from said  
26 subset are humanized.

1           Claim 22. The method for treating a patient suffering  
2   from a cancerous disease in accordance with claim 12  
3   wherein:

4           said method of production utilizes a tissue sample  
5   containing cancerous and non-cancerous cells obtained from  
6   a particular individual.

7  
8  
9  
10          Claim 23. Anti-cancer antibodies or fragments thereof  
11   selected from the group consisting of a 1LN-8, 4BD-1, a  
12   4BD-3, a 4BD-6, a 4BD-9, a 4BD-13, a 4BD-18, a 4BD-20, a  
13   4BD-25, a 4BD-26, a 4BD-27, a 4BD-28, a 4BD-32, a 4BD-37,  
14   a 4BD-50, a 6BD-1, a 6BD-3, a 6BD-5, a 6BD-11, a 6BD-25, a  
15   7BD-7, a 7BD-12-1, a 7BD-12-2, a 7BD-13, a 7BD-14, a 7BD-  
16   19, a 7BD-21, a 7BD-24, a 7BD-29, a 7BD-30, a 7BD-31, a  
17   7BDI-17, a 7BDI-58, a 7BDI-60, a 7BDI-62, a 5LAC2, a  
18   5LAC4, a 5LAC20, a 5LAC23, a H460-1, a H460-4, a H460-5, a  
19   H460-10, a H460-14, a H460-16-1, a H460-16-2, a H460-22-1,  
20   a H460-23 and a H460-27 monoclonal antibody or combinations  
21   thereof.

22  
23          Claim 24. Anti-cancer antibodies or fragments thereof  
24   produced by a hybridoma cell line having an ATCC Accession  
25   Number selected from the group  
26   consisting of           (to be provided before publication).

27

28

1           Claim 25. The use of a composition for treating a  
2   patient suffering from a cancerous disease by  
3   administration of an effective amount of said composition  
4   to a patient to mediate treatment of said cancerous  
5   disease, wherein said composition comprises one or more  
6   antibodies or fragments thereof selected from a subset of  
7   said antibodies or fragments in admixture with a  
8   pharmaceutically acceptable adjuvant, said anti-cancer  
9   antibodies or fragments thereof produced in accordance  
10   with a method for the production of individually  
11   customized anti-cancer antibodies which are useful in  
12   treating cancerous disease, said subset of antibodies or  
13   fragments thereof characterized as being cytotoxic against  
14   cells of a cancerous tissue, as being essentially benign  
15   to non-cancerous cells and being selected from the group  
16   consisting of a 1LN-8, 4BD-1, a 4BD-3, a 4BD-6, a 4BD-9, a  
17   4BD-13, a 4BD-18, a 4BD-20, a 4BD-25, a 4BD-26, a 4BD-27,  
18   a 4BD-28, a 4BD-32, a 4BD-37, a 4BD-50, a 6BD-1, a 6BD-3,  
19   a 6BD-5, a 6BD-11, a 6BD-25, a 7BD-7, a 7BD-12-1, a 7BD-  
20   12-2, a 7BD-13, a 7BD-14, a 7BD-19, a 7BD-21, a 7BD-24, a  
21   7BD-29, a 7BD-30, a 7BD-31, a 7BDI-17, a 7BDI-58, a 7BDI-  
22   60, a 7BDI-62, a 5LAC2, a 5LAC4, a 5LAC20, a 5LAC23, a  
23   H460-1, a H460-4, a H460-5, a H460-10, a H460-14, a H460-  
24   16-1, a H460-16-2, a H460-22-1, a H460-23 and a H460-27  
25   monoclonal antibody or combinations thereof.

26

27

**STATEMENT UNDER ARTICLE 19 (1)**

The amendments include:

(1) amendments to claim 1 to correct editorial errors.

(2) amendments to claims 1, 23 and 25 to insert reference to "a H460-22-1," in the three claims, support for this appearing in Table 5 on page 37, line 13.

To provide consistency with the above, a new disclosure page 36 is included with the amended claim pages.

Finally, attached hereto are copies of submissions and acknowledgement of the submission of Deposited Material and certain ATCC Numbers in relation to a related and counterpart U.S. application number 09/727,361 as provided to us by the instructing U.S. Attorney. This material also makes reference to H460-22-1.

The cell line referred to in the ATCC material is found in the disclosure at the noted pages hereinbelow:

<u>Cell Line</u>	<u>Reference Pages</u>
1LN-8	22 (23)
5LAC20	33 (33)
3BD-26	19 (20)
3BD-8	19 (20)
7BD-14	27 (30)
3BD-27	19 (20)
H460-27	36 (37)
H460-23	36 (37)
H460-16-2	36 (37)
H460-22-1	36 (37) (amended as above)
7BDI-60	27 (31)

The first reference page number being reference to the page where the cell line is listed, whereas the second bracketed number is the page of the table in which the cell line appears.

Appropriate amendments to the disclosure to add the ATCC numbers will be effected in due course.



## IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

IN RE APPLICANT : Young et al.  
 INVENTION : Individualized Anti-Cancer Antibodies  
 SERIAL NUMBER : 09/727,361  
 FILING DATE : November 29, 2000  
 EXAMINER : Susan Ungar  
 GROUP ART UNIT : 1642  
 OUR FILE NO. : 2056.009

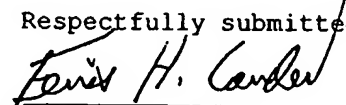
-----

To: The Commissioner of Patents and Trademarks  
 Washington, D.C. 20231

CORROBORATION FOR DEPOSITED MATERIALS

I, Ferris H. Lander, a person in a position to corroborate the identity of the hybridoma cell lines 1LN-8 (shown in the table on page 30), 3BD-8 (shown in the table on page 26), 3BD-26 (shown in the table on page 26), 3BD-27 (shown in the table on page 26), H460-27 (shown in the table on page 45), H460-23 (shown in the table on page 45), 7BD-14 (shown in the table on page 35) and 5LAC20 (shown in the table on page 41) which were deposited, in accordance with the Budapest Treaty, with the American Type Culture Collection (ATCC), 10801 University Blvd., Manassas, VA 20110-2209 on November 21, 2000 under Accession Numbers PTA-2693, PTA-2696, PTA-2695, PTA-2698, PTA-2699, PTA-2700, PTA-2697 and PTA 2694 respectively. And additionally, the hybridoma cell lines H460-16-2 (shown in the table on page 45) and 7BDI-60 (shown in the table on page 39) which were deposited on September 4, 2002 under Accession Numbers PTA-4621 and PTA-4623 respectively, do hereby state that the deposited hybridoma are the same hybridoma cell lines disclosed and claimed in the above-referenced patent application.

Respectfully submitted,

  
 Ferris H. Lander  
 Registration # 43,377

McHale & Slavin, P.A.  
 4440 PGA Blvd., Suite 402  
 Palm Beach Gardens, FL 33410  
 (561) 625-6575 (Voice)  
 (561) 625-6572 (Fax)

# ATCC

10801 University Blvd • Manassas, VA 20110-2209 • Telephone: 703-365-2700 • FAX: 703-365-2745

## BUDAPEST TREATY ON THE INTERNATIONAL RECOGNITION OF THE DEPOSIT OF MICROORGANISMS FOR THE PURPOSES OF PATENT PROCEDURE

### INTERNATIONAL FORM

RECEIPT IN THE CASE OF AN ORIGINAL DEPOSIT ISSUED PURSUANT TO RULE 7.3  
AND VIABILITY STATEMENT ISSUED PURSUANT TO RULE 10.2

To: (Name and Address of Depositor or Attorney)

Arius Research Inc.  
Attn: Lisa Cecchetto  
6299 Airport Road  
Mississauga, ON  
L4V 1N3  
Canada

Deposited on Behalf of: Arius Research Inc.

#### Identification Reference by Depositor:

Mouse hybridoma cell line: 1LN8  
Mouse hybridoma cell line: 5LAC20  
Mouse hybridoma cell line: 3BD26  
Mouse hybridoma cell line: 3BD8  
Mouse hybridoma cell line: 7BD14  
Mouse hybridoma cell line: 3BD27  
Mouse hybridoma cell line: H460-27  
Mouse hybridoma cell line: H460-23

#### Patent Deposit Designation

PTA-2693  
PTA-2694  
PTA-2695  
PTA-2696  
PTA-2697  
PTA-2698  
PTA-2699  
PTA-2700

The deposits were accompanied by:    a scientific description    a proposed taxonomic description indicated above. The deposits were received November 21, 2000 by this International Depository Authority and have been accepted.

AT YOUR REQUEST:   X   We will inform you of requests for the strains for 30 years.

The strains will be made available if a patent office signatory to the Budapest Treaty certifies one's right to receive, or if a U.S. Patent is issued citing the strains, and ATCC is instructed by the United States Patent & Trademark Office or the depositor to release said strains.

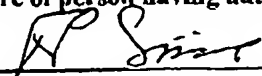
If the cultures should die or be destroyed during the effective term of the deposit, it shall be your responsibility to replace them with living cultures of the same.

The strains will be maintained for a period of at least 30 years from date of deposit, or five years after the most recent request for a sample, whichever is longer. The United States and many other countries are signatory to the Budapest Treaty.

The viability of the cultures cited above was tested November 30, 2000. On that date, the cultures were viable.

International Depository Authority: American Type Culture Collection, Manassas, VA 20110-2209 USA.

Signature of person having authority to represent ATCC:

  
Frank Simone, Director, Patent Depository

Date: December 21, 2000

cc: Mr. Ferris Lander (Ref: Docket or Case No.: 09/415,278)

# ATCC

10801 University Blvd • Manassas, VA 20110-2209 • Telephone: 703-365-2700 • FAX: 703-365-2745

**BUDAPEST TREATY ON THE INTERNATIONAL RECOGNITION OF  
THE DEPOSIT OF MICROORGANISMS FOR THE PURPOSES OF PATENT PROCEDURE**

**INTERNATIONAL FORM**

**RECEIPT IN THE CASE OF AN ORIGINAL DEPOSIT ISSUED PURSUANT TO RULE 7.3  
AND VIABILITY STATEMENT ISSUED PURSUANT TO RULE 10.**

To: (Name and Address of Depositor or Attorney)

Arius Research Inc.  
Attn: Jean de Sousa-Hitzler  
55 York Street 16<sup>th</sup> Floor  
Toronto, Ontario  
Canada M5J 1R7

Deposited on Behalf of: Arius Research Inc.

Identification Reference by Depositor:

Patent Deposit Designation

Mouse Hybridoma: H460-16-2  
Mouse Hybridoma: H460-22-1  
Mouse Hybridoma: 7BD1-60

PTA-4621  
PTA-4622  
PTA-4623

The deposits were accompanied by:    a scientific description    a proposed taxonomic description indicated above. The deposits were received September 4, 2002 by this International Depository Authority and have been accepted.

AT YOUR REQUEST:   X   We will inform you of requests for the strains for 30 years.

The strains will be made available if a patent office signatory to the Budapest Treaty certifies one's right to receive, or if a U.S. Patent is issued citing the strains, and ATCC is instructed by the United States Patent & Trademark Office or the depositor to release said strains.

If the cultures should die or be destroyed during the effective term of the deposit, it shall be your responsibility to replace them with living cultures of the same.

The strains will be maintained for a period of at least 30 years from date of deposit, or five years after the most recent request for a sample, whichever is longer. The United States and many other countries are signatory to the Budapest Treaty.

The viability of the cultures cited above was tested September 6, 2002. On that date, the cultures were viable.

International Depository Authority: American Type Culture Collection, Manassas, VA 20110-2209 USA.

Signature of person having authority to represent ATCC:

Marie Harris  
Marie Harris, Patent Specialist, ATCC Patent Depository

Date: October 9, 2002

cc: Mr. Ferris Lander  
(Ref: Docket or Case No.: 2056.009 & US Serial No. 09/727361)

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## INTERNATIONAL SEARCH REPORT

tional Application No

PCT/CA 01/01838

## A. CLASSIFICATION OF SUBJECT MATTER

IPC 7 A61K39/395 A61K47/48 A61P35/00

According to International Patent Classification (IPC) or to both national classification and IPC

## B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

IPC 7 G01N C07K

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

MEDLINE, BIOSIS, EPO-Internal, WPI Data, PAJ

## C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	US 2001/009665 A1 (YOUNG DAVID S F ET AL) 26 July 2001 (2001-07-26) the whole document ---	1-26
X	US 2001/003777 A1 (YOUNG DAVID S F ET AL) 14 June 2001 (2001-06-14) the whole document ---	1-26
X	US 6 180 357 B1 (YOUNG DAVID S F ET AL) 30 January 2001 (2001-01-30) the whole document --- -/--	1-26



Further documents are listed in the continuation of box C.



Patent family members are listed in annex.

## \* Special categories of cited documents:

- \*A\* document defining the general state of the art which is not considered to be of particular relevance
- \*E\* earlier document but published on or after the international filing date
- \*L\* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
- \*O\* document referring to an oral disclosure, use, exhibition or other means
- \*P\* document published prior to the international filing date but later than the priority date claimed

- \*T\* later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
- \*X\* document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
- \*Y\* document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.
- \*G\* document member of the same patent family

Date of the actual completion of the international search

10 January 2003

Date of mailing of the international search report

16/01/2003

Name and mailing address of the ISA

European Patent Office, P.B. 5818 Patentlaan 2  
NL - 2280 HV Rijswijk  
Tel. (+31-70) 340-2040, Tx. 31 651 epo nl,  
Fax: (+31-70) 340-3016

Authorized officer

COVONE-VAN HEES, M

## INTERNATIONAL SEARCH REPORT

International Application No.

PCT/CA 01/01838

## C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	<p>WO 95 20401 A (UNIV BOSTON) 3 August 1995 (1995-08-03)</p> <p>page 17, line 16-27 page 29, line 4-20 example 13 claims 30,31,38-40</p> <p>-----</p>	<p>1-6, 11-17, 22-26</p>

# INTERNATIONAL SEARCH REPORT

International application No.  
PCT/CA 01/01838

## Box I Observations where certain claims were found unsearchable (Continuation of Item 1 of first sheet)

This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1. ☒ Claims Nos.:  
because they relate to subject matter not required to be searched by this Authority, namely:  
  
Although claims 1-22, 25,26 are directed to a method of treatment of the human/animal body, the search has been carried out and based on the alleged effects of the compound/composition.
2. ☐ Claims Nos.:  
because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:
3. ☐ Claims Nos.:  
because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).

## Box II Observations where unity of invention is lacking (Continuation of Item 2 of first sheet)

This International Searching Authority found multiple inventions in this International application, as follows:

1. ☐ As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2. ☐ As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. ☐ As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4. ☐ No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:

Remark on Protest

- ☐ The additional search fees were accompanied by the applicant's protest.
- ☐ No protest accompanied the payment of additional search fees.

Patent document cited in search report		Publication date	Patent family member(s)	Publication date
US 2001009665	A1	26-07-2001	US 6180357 B1	30-01-2001
			US 2002041877 A1	11-04-2002
			US 2001003777 A1	14-06-2001
<hr/>				
US 2001003777	A1	14-06-2001	US 6180357 B1	30-01-2001
			US 2002041877 A1	11-04-2002
			US 2001009665 A1	26-07-2001
<hr/>				
US 6180357	B1	30-01-2001	US 2002041877 A1	11-04-2002
			US 2001009665 A1	26-07-2001
			US 2001003777 A1	14-06-2001
<hr/>				
WO 9520401	A	03-08-1995	AU 1736495 A	15-08-1995
			EP 1231268 A2	14-08-2002
			EP 0744958 A1	04-12-1996
			US 6335163 B1	01-01-2002
			WO 9520401 A1	03-08-1995
			US 5789208 A	04-08-1998
			US 2001049107 A1	06-12-2001
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